

Method and Nucleic Acids for the Detection of Microorganisms Relevant to Brewing

The invention relates to a method for the detection of microorganisms relevant to brewing, as well as to nucleic acids and combinations thereof which can be used in this method. The invention further relates to the use of the nucleic acids according to the invention or combinations thereof for the detection and/or for the identification and/or characterisation of different genera or species of microorganisms relevant to brewing.

Beer can be regarded as very stable microbiologically, and can only be spoilt by a relatively manageable number of bacteria. In order to discover contamination with these organisms as early as possible, an analytical system which allows rapid detection of the microorganisms in the matrix beer must be used, since countermeasures must be undertaken immediately.

The common feature of all microorganisms harmful to beer is the trace contamination of individual vessels (barrels, bottles) and their slow growth. In particular, microbiological culturing of the anaerobic microorganisms is very difficult. The beer-spoiling bacteria at present known are classed into the following genera: *Lactobacillus, Pediococcus, Pectinatus* and *Megasphaera*. Members of the *Selenomonas* and *Zymophilus* genera have not yet emerged as beer contaminants; however, contamination of beer and their subsequent growth in it cannot be ruled out.

The genus *Lactobacillus* describes Gram positive, non-sporulating, mostly immotile and chain-forming rods, which are long, thin and sometimes curved. Coccoid forms are also sometimes observed. Members of the genus *Lactobacillus* are microaerophilic, and some are anaerobic. They are cytochrome- and catalase-negative, their metabolism is fermentative and they require a complex nutrient medium. The molar G+C content of the DNA is between 32 and 53%.

As well as in beer, *Lactobacilli* are found in dairy and cereal products, in meat and fish products, in water, waste water, wine, fruit and fruit juices, acid-pickled vegetables, sauerkraut, silage and sourdough. Although they are a component of the normal oral, intestinal and vaginal flora of mammals, they are however seldom pathogenic (*Bergeys Manual of Syst. Microbiology*, 1984, p. 1209-1234). In beer, because of their metabolic

products, they lead to clouding and undesired flavour changes. Species relevant to beer spoilage are Lactobacillus brevis, Lactobacillus lindneri, Lactobacillus casei, Lactobacillus paracasei, Lactobacillus coryniformis and Lactobacillus curvatus (Back, Brauwelt, 1980, 120, p. 1562-1569).

The genus *Pediococcus* includes Gram positive, immotile and non-sporulating cocci. They form tetrads or occur as pairs. They are facultative anaerobes, and their oxygen sensitivity differs from species to species. *Pediococci* are cytochrome and catalasenegative and require a complex nutrient medium (*Bergeys Manual of Syst. Microbiology*, 1984, p. 1075-1079). They are used as starter cultures for the production of raw sausage products, they ferment various types of pickled vegetables and lead to the spoilage of foodstuffs (Firnhaber, Baumgart: *Mikrobiologische Untersuchung von Lebensmitteln*, 1993, p. 413-419, 115-117). The genus includes 8 species, and the species *Pediococcus damnosus* and *Pediococcus inopinatus* should be regarded as harmful to beer.

The genus *Pectinatus* includes the species *Pectinatus cerevisiiphilus*, *Pectinatus frisingiensis* and the strain *Pectinatus sp.* DSM 20764, not further taxonomically classified. All strains have been isolated from spoilt beer (Schleifer et al., *Int. J. of Syst. Bacteriology*, 1990, p. 19-27). These are slightly bent, non-sporulating rod-shaped bacteria. They have comb-like flagella, and are motile. They produce neither catalase nor cytochrome oxidase, and are obligate anaerobes. The molar G+C content is 38-41%. In the genus *Pectinatus*, and also in the genera *Megasphaera*, *Selenomonas* and *Zymophilus*, the cell wall is more similar to the Gram-positive bacteria than to the Gram-negative bacteria. Although the Gram staining is negative, they are taxonomically classified among the Gram-positive bacteria (Haikara, *The Prokaryotes*, 2nd Edition, Vol. II, 1991, p. 1993-2004).

The genus *Megasphaera* includes the species *Megasphaera elsdenii* and *Megasphaera cerevisiae*. Only *Megasphaera cerevisiae* is relevant to brewing, and is described as a Gram negative, strictly anaerobic, cytochrome- and catalase-negative, immotile and sometimes slightly stretched coccus, which occurs singly, in pairs or in short chains. The mean cell diameter is about 1.4 μ m, and the molar G+C content 42.4-44.8%. Main metabolites are sulphur compounds, such as H₂S and volatile fatty acids. In beer,

contamination with *Megasphaera cerevisiae* leads to very marked changes in aroma and taste (Haikara, *The Prokaryotes*, 2nd Edition, Vol. II, 1991, p. 1993-2004).

Species of the genus *Selenomonas* are defined as obligate anaerobes, Gram negative, non-sporulating, slightly curved and motile rods. The molar G+C content is about 48-58% (Schleifer et al., *Int. J. of Syst. Bacteriology*, 1990, p. 19-27). *Selenomonads* are isolated from the stomach and intestinal tract and the dung of mammals. The genus includes 10 species (Hespell et al., *The Prokaryotes*, 2nd Edition, Vol. II, 1991, p. 2005-2013). Only *Selenomonas lacticifex* has been isolated from starter yeast, and is thus relevant to brewing. *Selenomonas lacticifex* has not yet emerged as a beer-spoiling bacterium; however, its growth in beer is possible, and hence it fulfils the definition of a beer-spoiling organism.

The species *Zymophilus paucivorans* and *raffinosivorans* belong to the genus *Zymophilus* as Gram-negative, slightly bent, motile rods, which occur singly, in pairs or in short chains. The molar G+C content is about 38-41%. They are obligate anaerobes and have a fermentative metabolism. Both species are isolated from starter yeasts and brewery wastes; growth in beer has only been observed with *Zymophilus raffinosivorans* (Schleifer et al., *Int. J. of Syst. Bacteriology*, 1990, p. 19-27).

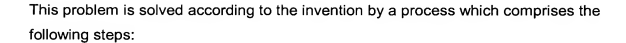
On the basis of comparison of the 16S rRNA gene sequences, all the genera to be tested are classified among the Gram-positive bacteria with low G+C content. The genera *Pediococcus* and *Lactobacillus* are classified into the *Lactobacillaceae* family, and the genera *Pectinatus*, *Megasphaera*, *Selenomonas* and *Zymophilus* into the *Sporomusa* group. The *Sporomusa* group is also described as a group of the Grampositive Eubacteriales with Gram-negative cell wall (Stackebrandt et al., *The Prokaryotes*, 2nd Edition, Vol. II, 1991, p. 25-26, 33).

A classical microbiological determination of the microorganisms described above can require up to 10 days. However, a markedly faster analysis is desirable, as otherwise unnecessary storage costs arise or the beer being tested has already been delivered. For these reasons, several rapid detection methods have already been developed. Thus, for example, organisms harmful to beer can be detected on the basis of their metabolic products (Haikara et al. Microbiology, 1995, 141, p. 1131-1137). Other indirect methods are turbidometry (Haikara et al., *ASBC*, 1990, p. 92-95) and measurement of

the ATP bioluminescence (Miller et al., *J. Inst. Brew.*, 1989, Vol. 95, p. 317-319). Detection by means of antibodies is also rapid and specific (Gares et al., *ASBC*, 1993, p. 158-163; Winnewisser et al., *Int. J. of Bacteriology*, 1995, 45, p. 403-405). With these methods, the disadvantage is that either non-specific parameters are tested or only one species or genus is detected in each case. Also, the equipment and staff cost is high. An overview of rapid methods for the detection of contaminants relevant to brewing is given by Dowhanick (*Cerevisia*, 1995, 20/4, p. 40-49).

The polymerase chain reaction (PCR; Mullis et al., see US 4,683,195, US 4,683,202 and US 4,965,188) is a rapid and effective method of specifically detecting organisms. A range of nucleic acids are known, through the use of which as primers and/or probes the specific detection of microorganisms relevant to brewing is possible. However, a disadvantage is that with the use of these nucleic acid molecules in an amplification or detection reaction, it is always only possible to detect a fraction of all possible microorganisms relevant to brewing. These PCR systems serve for the specific detection in each case only of individual species in an amplification reaction of the genera Lactobacillus, Pediococcus, Pectinatus and Mega-sphaera (Sakamoto US 5,869,642; Nietupski et al., US 5,705,339 and US 5,484,900; Tsuchia et al., JP 06141899A, JP 06113888A / *ASBC J.*, 1992, p. 64-67 / *ASBC J.*, 1993, p. 40-41; Yasui JP07289295A / *Can. J. Microbiol.*, 1997, 43, p. 157-163, Shimada et al., JP06090793; Alatossava et al. WO97/09448; Doyle et al., J. of Ind. Microbiology, 1995, 15, p. 67-70; DiMichele et al., ASBC J., 1993, p. 63-66; Vogeser et al, Brauwelt, 1998, 24/25, p. 1060-1063). Further, the methods described for visualisation of the amplification products, such as, for example, agarose gel electrophoresis, present problems, as the carcinogenic and highly toxic ethidium bromide is used for staining the amplification products. These methods can only be automated with difficulty and the assessment of the agarose gels or the identification of the microorganisms on the basis of the length of the amplification products is sometimes not clear.

The problem to be solved by the present invention was, therefore, to provide a method and means which make possible a rapid test of beer and brewing raw materials for contamination with microorganisms, the test being required to detect the whole range of possible beer-contaminating microorganisms.



- (a) bringing the sample into contact with a combination of at least two first nucleic acid molecules (primers), which hybridise with a region of a microbial nucleic acid conserved in microorganisms relevant to brewing;
- (b) amplification of the microbial nucleic acid or a portion thereof to produce at least one amplification fragment;
- (c) bringing the amplification fragments obtained in step (b) into contact with at least one second nucleic acid molecule (probe), which specifically hybridises with at least one amplification fragment that comprises a sequence of the microbial nucleic acid specific for all microorganisms relevant to brewing or for one or several families, genera or species of microorganisms relevant to brewing; and
- (d) detection of at least one hybrid nucleic acid which consists of an amplification fragment and a second nucleic acid molecule introduced in step (c),

and by a nucleic acid molecule selected from:

- (i) a nucleic acid with a sequence according to SEQ ID NO 1-107 or a fragment thereof at least 10, preferably 15-30, nucleotides long;
- (ii) a nucleic acid which specifically hybridises with a nucleic acid according to (i);
- (iii) a nucleic acid which is at least 70%, preferably at least 90%, identical with a nucleic acid according to (i) or (ii), or
- (iv) a nucleic acid which is complementary to a nucleic acid according to (i) to (iii).

In the sequences according to SEQ ID NO 1-107, nucleotides are abbreviated as follows: G = guanosine, A = adenosine, T = thymidine, C = cytidine, U = uracil, i = inosine. In accordance with IUPAC, mixtures are abbreviated as follows: R = G or A, Y = C or A, A or A

For the determination of identity (in the sense of complete agreement, corresponding to 100% identity) with nucleic acid sequences according to (iii), partial sequences of a larger polynucleotide are considered. These partial sequences include 10 nucleotides and are identical when all 10 building blocks are identical in the two sequences compared. The nucleotides thymidine and uridine are to be regarded as identical. All possible fragments of a larger polynucleotide can be regarded as partial sequences.

Here 90% identity is present, when in the two sequences to be compared 9 out of 10 or 18 out of 20 nucleotides in one section are identical.

As an example, let us consider two polynucleotides which comprise 20 nucleotides and differ in the 5th element. In a sequence comparison, six 10-nucleotide ones are then found which are identical, and 5 which are not identical, as they differ in one element.

Otherwise, the identity can also be determined by degree, the unit being stated in percent. For determination of the degree of identity, partial sequences are also considered, which as a minimum include the length of the sequence actually used, e.g., as primer, or else 20 nucleotides.

As an example, polynucleotides A with a length of 100 nucleotides and B with a length of 200 nucleotides are compared. From polynucleotide B, a primer with a length of 14 nucleotides is derived. For the determination of the degree of identity, polynucleotide A is compared with the primer over its whole length. If the sequence of the primer occurs in polynucleotide A, but differs in one element, then there is a fragment with a degree of identity of 13/14 → 92.3%.

In the second example, the whole of the aforesaid polynucleotides A and B are compared. In this case, all possible comparison windows of a length of 20 nucleotides are applied, and the degree of identity determined for them. Thus, if nucleotides 50-69 of polynucleotide A and B are identical with the exception of nucleotide No. 55, then for these fragments a degree of identity of 19/20 → 95% is found.

The method according to the invention can be carried out more rapidly than the previous microbiological detection methods, and makes it possible to detect several, preferably all, microorganisms relevant to brewing potentially present in a sample, such as, for example, even *Lactobacillus* species or members of the genera *Selenomonas* or

Zymophilus seldom arising as contaminants, for which hitherto no detection method existed. The detection is comprehensive and indicates all contamination risks in the brewery. By means of the method according to the invention, microorganisms relevant to brewing can be detected both in beer samples and also in raw material samples (barley malt, yeast, hops, water) or samples of intermediate products in beer production (e.g. mash, wort) even when the number of contaminating microorganisms is still low.

In this context, microorganisms relevant to brewing are understood primarily to mean bacteria and in particular the bacteria described above, Lactobacillus brevis, Lactobacillus lindneri, Lactobacillus casei, Lactobacillus paracasei, Lactobacillus coryniformis, Lacto-bacillus curvatus, Pediococcus damnosus, Pediococcus inopinatus, Pectinatus cerevisii-philus, Pectinatus frisingiensis, Pectinatus sp. DSM 20764, Megasphaera cerevisiae, Selenomonas lacticifex, Zymophilus paucivorans and Zymophilus raffinosivorans, and also all microorganisms to be found in beer, which, while they do not belong to the aforesaid species, can nonetheless multiply in beer, for example, rare members of the Lactobacillaceae family, such as Lactobacillus malefermentans, Lactobacillus buchneri, Lactobacillus parabuchneri, Lactobacillus sanfrancisco, Lactobacillus delbrueckii, Leuconostoc mesenteroides, Pediococcus pentosacaeus and Lactococcus lactis.

The microorganisms detectable by the method according to the invention are, thus, not limited to the microorganisms hitherto described as beer contaminants. Rather, the use of the nucleic acid molecules and the method according to the invention offers the possibility of recognising the presence of other microorganisms relevant to brewing, which have not previously been described as beer contaminants. A positive result at the level of higher taxonomic units (e.g. orders, families, genera) combined with a negative result at the level of the lower taxonomic units known to be relevant to brewing (e.g. species, subspecies, strains) indicates a contamination with such a non-typical microorganism relevant to brewing.

In a first step of the method according to the invention, the sample to be tested is brought into contact with a combination of at least two first nucleic acid molecules (primers). These nucleic acid molecules hybridise with a region of a microbial nucleic acid which is conserved in microorganisms relevant to brewing. The hybridisation takes place through pairing of the primer with regions of the microbial nucleic acid which have

an at least partly complementary base sequence. The term "conserved" characterises the evolutionary variability of nucleotide sequences for species of different taxonomic units. If corresponding sequence sections from at least two microorganisms relevant to brewing are compared, the sequence can be regarded as variable or as conserved. Comparison sequences which are at least 95% identical are described as conserved, and those which are less than 95% identical as variable. Thus, a region of a nucleic acid conserved in microorganisms relevant to brewing denotes a region which is at least 95% identical in all microorganisms relevant to brewing (as defined above).

In a preferred embodiment of the present invention, the conserved region occurs in a genome section which contains the bacterial 23S and 5S genes. This region includes the intergenic spacer between the genes for the 23S rRNA and the 5S rRNA and the bounding 23S and 5S rDNA genes, and includes both conserved sequence regions and also hypervariable (i.e., very organism-specific) sequence regions. Prokaryotic ribosomes as a rule contain three distinct nucleic acid components, which are generally known as 5S, 16S and 23S rRNA (ribosomal nucleic acid). The genetic information for these ribonucleic acids (rDNA) is typically arranged in the genome as a tandem. The typical organisation of such a unit is 16S-23S-5S, where the genes are connected to one another by short hypervariable intergenic regions, so-called spacers. The units are present several times in the genome, and the number of operons can vary from species to species. The high conservation of the DNA sequence in certain sections of the ribosomal DNA over the whole bacterial kingdom allows the design of non-specific oligonucleotides even without exact knowledge of the individual DNA sequences of the organisms to be investigated. The sequences according to SEQ ID NO 1-20 according to the invention (Table 1) are sequences of the 23S-5S intergenic spacer of microorganisms relevant to brewing, from which nucleic acid molecules for use in the method according to the invention can be derived.

The combination of at least two first nucleic acid molecules used in the first step of the method according to the invention is selected, such that they are usable as primers in an amplification reaction, i.e., one nucleic acid molecule hybridises onto a first conserved region of the first strand of the target DNA and the other nucleic acid onto a second conserved region of the DNA strand complementary to the first, wherein the desired target region of the DNA is included. Both nucleic acid molecules have a length of at

least 10 bp, preferably 15-30 bp. In a preferred embodiment of the invention, a combination of at least two nucleic acid molecules according to this invention is used. In a particularly preferred embodiment of the invention, a combination is used which includes at least one nucleic acid molecule with a sequence according to one of the SEQ ID NO 40 to 47 (Table 2) and at least one nucleic acid molecule with a sequence according to SEQ ID NO 48-54 or SEQ ID NO 55-59 or SEQ ID NO 60-72 (Table 2).

In a second step of the method according to the invention, the microbial nucleic acid or a portion thereof is amplified, whereby at least one amplification fragment is produced. Amplification is understood to mean the raising of the concentration of a nucleic acid or a portion thereof present in a reaction mixture. Processes used for the amplification of nucleic acids are for example the PCR (US 4,683,195, US 4,683,202 and US 4,965,188), the "self-sustained sequence replication" (EP 329,822), the "transcription-based amplification system" (EP 310,229) and the "β-RNA replicase system" (US 4,956,858). In a preferred embodiment of the present invention, the amplification comprises a polymerase chain reaction (PCR). In a further embodiment of the present invention, the amplification comprises a ligase-chain reaction or an isothermal nucleic acid amplification.

In a third step of the method according to the present invention, the amplification fragments obtained are brought into contact with at least one second nucleic acid molecule (probe). This nucleic acid molecule or these nucleic acid molecules hybridise specifically with at least one amplification fragment that comprises a sequence of the microbial nucleic acid which is specific for all microorganisms relevant to brewing or for one or several families, genera or species of microorganisms relevant to brewing, i.e., only occurs in members of these families or genera or in these species.

The double-strand formation of two identical or similar nucleotide fragments (DNA, RNA, PNA) is described as hybridisation. The term specific hybridisation is used when a stable hybrid nucleic acid between the oligonucleotide and the corresponding target DNA of the oligonucleotide exists, but not to other DNA than the target DNA. For the purposes of this invention, the feature "sequence which specifically hybridises with a sequence according to (i)" refers to a sequence, which under stringent conditions, hybridises with the sequence according to (i). For example, the hybridisations can be carried out at 50°C with a hybridisation solution consisting of 2.5 x SSC, 2 x Denhardts solution, 10 mM

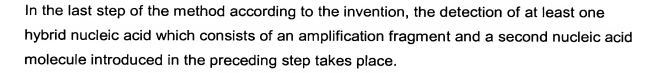
Tris, 1 mM EDTA pH 7.5. Suitable washing conditions are for example four times repeated 1-minute washings in 0.1 x SSC to 1.0 x SSC, 2 x Denhardts, 10 mM Tris, 1 mM EDTA, pH 7.5 at 20-50°C.

In a preferred embodiment of the invention, one or several of the nucleic acid molecules according to the invention is used as a second nucleic acid molecule (probe). Consensus probe is understood to mean a nucleic acid molecule which hybridises with highly conserved regions of a microbial nucleic acid and reacts with the amplification products of all microorganisms relevant to brewing. Nucleic acid molecules according to the invention which are usable as consensus probes have a sequence according to one of SEQ ID NO 40 to 72 (Table 2).

For the detection of a specific genus of microorganisms relevant to brewing, a nucleic acid molecule with a sequence according to one of SEQ ID NO 35 to 39 or SEQ ID NO 104 to 107 (Table 2) is preferably used. The genus specificity of a probe is defined as the ability of this probe to hybridise with the DNA of all isolates of as large as possible a group of members of the particular genus to be detected.

Species-specific nucleic acid probes are understood to mean nucleic acid molecules which hybridise with the DNA of all isolates of the particular species to be detected under the same stringency conditions. Species-specific nucleic acid molecules according to the invention with SEQ ID NO 21-22, SEQ ID NO 25-34, SEQ ID NO 73-78, SEQ ID NO 80-85 or SEQ ID NO 87-97 (Table 2) can be used.

The probes SEQ ID NO 23-24, SEQ ID NO 79, SEQ ID NO 86 and SEQ ID NO 98 to 103 are special cases. With the probes according to SEQ ID NO 23 and SEQ ID NO 79, strains of *Lactobacillus casei* and *Lactobacillus paracasei* ssp. *paracasei* can be detected. A probe according to SEQ ID NO 24 allows the detection of two subspecies of *Lactobacillus coryniformis* (*L. coryniformis* ssp. *coryniformis* and *L. coryniformis* ssp. *torquens*). With the probe SEQ ID NO 86, strains of the species *Pediococcus damnosus*, *Pediococcus inopinatus* and *Pediococcus parvulus* can be detected. With the use of these probes, other microorganisms relevant to brewing are not detected. Likewise, with the probes SEQ ID NO 98 to 103, all species of the *Lactobacillaceae* family relevant to brewing to be detected are detected, and other species and genera relevant to brewing are discriminated against.

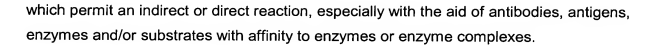


Preferably, first nucleic acid molecules (primers) and/or second nucleic acid molecules (probes) are at least 10 nucleotides, preferably 15-30 nucleotides long. In one embodiment of the present invention, the first and/or the second nucleic acid molecules are modified in that up to 20% of the nucleotides in 10 consecutive nucleotides, in particular 1 or 2 nucleotides of a block of 10 are replaced by nucleotides which do not occur naturally in bacteria.

The method according to the invention preferably includes the so-called consensus PCR. In this method, multiplication of the microbial nucleic acid or a portion thereof, and subsequent detection of these molecules by hybridisation with labelled specific probes take place. In the consensus PCR, nucleic acid molecules are used which make it possible to obtain an amplification product from several or, indeed, all of the relevant strains, subspecies, species or genera. The amplification does not lead to a differentiation of the microorganisms. The specificity of the detection is achieved through the subsequent hybridisation reaction with specific probes. In this way, microorganisms relevant to brewing can be simultaneously detected in a simple combination of amplification and detection reaction.

This kind of amplification and detection makes it possible to automate the detection reaction, so that a high sample throughput becomes possible. For example, a PCR-ELISA detection procedure can be used, in which the respective probes are bound in different wells of a microtitre plate, in which the hybridisation and the detection of the labelled amplification products then occurs. The detection can also be effected by the use of a microarray, on which several probes are immobilised, as a result of which the detection reaction can be carried out quickly and at no great cost.

In a preferred embodiment of the invention, the second nucleic acid molecule (probe) is modified or labelled in such a way that it can produce a detectable signal. The modification or labelling is selected from (i) radioactive groups, (ii) coloured groups, (iii) fluorescent groups, (iv) groups for immobilisation on a solid phase and (v) groups



For the purposes of this invention, labelling indicates directly or indirectly detectable groups or groups for immobilisation on a solid phase, which are attached to the nucleic acid molecule. Directly detectable are metal atoms, radioactive, coloured or fluorescent groups. Indirectly detectable are immunologically or enzymatically detectable groups, for example, antigens and antibodies, haptens or enzymes or enzymatically active parts of enzymes. These indirect groups are detected in subsequent reactions. Preferred are haptens which are coupled to an oligonucleotide and which are detected in a subsequent antibody reaction.

The nucleic acid molecules according to the invention can be used for the detection and/or for the identification and/or characterisation of bacteria relevant to brewing. The primers and/or probes described herein can also be used in the detection of the described microorganisms in drinks other than beer, in other samples from the brewing sector, such as for example in raw materials, starter yeast, environmental samples, in other foodstuff samples or in clinical samples, etc.

Examples:

Example 1: Determination of the DNA target sequence of the bacteria harmful to beer and closely related species

By sequence comparison of known 23S rDNA and 5S rDNA sequences (GenBank Sequence Database of the National Center of Biotechnology Information: NCBI), conserved gene regions were identified, which serve as hybridisation sites for the primers used for the sequencing. From pure cultures of the bacteria listed in Table 1, genomic DNA was isolated by known standard methods. With primers which hybridise in highly conserved regions, amplification products of all bacteria to be detected were obtained in a PCR. The following primers were used for the amplification and the subsequent sequencing:

Primer 1 = SEQ ID NO 47: 5'-AAG TGC TGA AAG CAT CTA AG-3' Primer 2 = SEQ ID NO 55: 5'-GGC RRY GTC TAY TYT CSC-3'

Composition of the PCR:

Genomic DNA (10 – 100 ng)	1.00 μΙ	
H ₂ O	16.85 μΙ	
Buffer (10 x)	2.50 μl	1 x
dNTP (10 mM)	0.50 μl	200 μΜ
Primer 1 = Seq ID NO 48 (5 μM)	1.50 μΙ	0.30 μΜ
Primer 2 = Seq ID NO 49 (5 μM)	1.50 μl	0.30 μΜ
MgCl ₂ (50 mM)	1.00 μΙ	2.00 mM
Taq-polymerase (5 U/μl)	0.15 μl	0.03 U/μl
Σ	25.00 μl	

Temperature profile:

5 mins	95°C	
30 secs	95°C	
30 secs	50°C	x 38
30 secs	72°C	
5 mins	72°C	

These amplification products were purified via an agarose gel and by a subsequent treatment with the QIAquick PCR Gel Extraction Kit (Quiagen Co.) and sequenced in the Long Read Sequencer Model 4000L (LI-COR Co.) with the aforesaid primers, which are provided with an IRD-800 label. The resulting sequences of the 23S/5S rDNA spacer regions of the bacteria relevant to brewing and the phylogenetically closely related species were compared with one another and sequence regions identified which:

- 1.) are to be found in all species of the particular genus to be detected and at the same time differ from those of other genera or species,
- 2.) are only to be found in the particular species to be detected, but differ from other bacteria to be detected and not to be detected.

In the sequence regions described under 1.), hybridisation sites of genus-specific oligonucleo-tides were defined, and in the sequence regions described under 2.), the binding sites of species-specific oligonucleotides were defined.

Example 2: Detection of Bacteria Harmful to Beer by the Polymerase Chain Reaction

I. Amplification

Genomic DNA was isolated from pure cultures of the bacteria listed in Table 1 by known standard methods. Decimal dilutions from 1 fg/ μ l to 1 pg/ μ l of these preparations were then used in a PCR with the following composition:

Primer 3 = SEQ ID NO 46:

5'-AAG GGC CAT CRC TCA ACG G -3'

Primer 4 = SEQ ID NO 48:

5'-TGT GTT CGi iAT GGG AAC AGG TG -3'

Genomic DNA	1.00 μl	4.00 μl	
H ₂ O	16.60 μl	66.40 μΙ	
Buffer (10 x)	2.50 μΙ	10.00 μl	1 x
dNTP (10 mM)	0.50 μΙ	2.00 μl	0.20 mM
Primer 3 = Seq ID NO 21 (5 μM)	1.50 µl	6.00 μl	0.30 mM
Primer 4 = Seq ID NO 22 (5 μM)	1.50 μl	6.00 μΙ	0.30 mM
digoxigenin labelled			
DMSO (100%)	0.25 μl	1.00 μl	1.00 %
MgCl ₂ (50 mM)	1.00 μl	4.00 μl	2.00 mM
Taq-polymerase (5 U/μl)	0.15 μl	0.60 μΙ	0.03 U/μl
Σ	25.00 μl	100.00 μl	

The PCR was performed under the following conditions in the Mastercycler® (Eppendorf Co.) according to the following temperature profile:

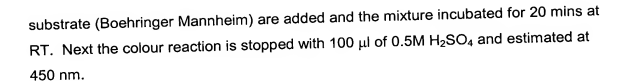
5 mins	95°C	
30 secs	95°C	
45 secs	55°C	x 38
90 secs	72°C	
5 mins	72°C	

Primer 3 (SEQ ID NO 46) was determined by sequence comparison of known 23S rDNA sequences (GenBank Sequence Database of NCBI). It hybridises onto highly conserved sequence sections in the 23S rDNA gene region. The binding site lies outside the region sequenced with the primers SEQ ID NO 48 and 49.

Primer 4 (SEQ ID NO 48) was determined on the basis of our own sequence data. The hybridisation site of primer 2 lies adjacent to the intergenic 23S/5S spacer in the 5S rDNA region.

II. Detection by PCR-ELISA

The detection is effected by PCR-ELISA. For this, per probe used, 5µl of amplification product are treated with 5 µl of denaturation buffer (125 mM NaOH, 20 mM EDTA, pH 14) and incubated for 15 mins at room temperature. Each time, 2 pmoles of the particular biotinylated probe are pipetted into 100 μl of hybridisation buffer (2.5 x SSC, 2 x Denhardts solution, 10 mM Tris, 1 mM EDTA, pH 7.5) and transferred to the wells of a microtitre plate coated with streptavidin and preincubated at the hybridisation temperature of 50°C. After the denaturation, the denaturation mixture is pipetted into the hybridisation mixture. Next the mixture is incubated for 30 minutes at hybridisation temperature. If the hybridisation is complete, the hybridisation mixture is removed and the plate washed 4x with 200 μ l of wash buffer 1 (WB1: 0.1 x SSC, 2 x Denhardts, 10 mM Tris, 1 mM EDTA, pH 7.6) for 1 min. each time at hybridisation temperature. Next, 100 μ l of a solution of a horseradish peroxidase conjugated anti-digoxigenin antibody diluted according to the manufacturer's instructions is added (Boehringer Mannheim). The conjugate is diluted in wash buffer 2 (WB2: 100 mM Tris, 150 mM NaCl, 0.05% Tween 20, 0.5% blocking reagent, 100 μg/ml herring sperm, pH 7.6). Next, the antibody incubation is performed at 37°C for 30 mins. After this, the plate is washed four times with 200 µl of WB2 (at room temperature). After the washing, 100 µl of POD



III. Assessment

According to the detection protocol described above, the detection was performed for all bacteria and bacteria groups investigated, using the corresponding genus- and speciesspecific probes. Genus-specific probes are SEQ ID NO 35 for Pediococcus, SEQ ID NO 36 for Pectinatus, SEQ ID NO 37 for Megasphaera, SEQ ID NO 38 for Selenomonas and SEQ ID NO 39 for Zymophilus. Species-specific probes are SEQ ID NO 21 for Lactobacillus brevis, SEQ ID NO 22 for Lactobacillus lindneri, SEQ ID NO 23 for Lactobacillus casei + paracasei, SEQ ID NO 24 for Lactobacillus coryniformis, SEQ ID NO 25 for Lactobacillus curvatus, SEQ ID NO 26 for Pediococcus damnosus, SEQ ID NO 27 for Pediococcus inopinatus, SEQ ID NO 28 for Pectinatus cervisiiphilus, SEQ ID NO 29 for Pectinatus frisingiensis, SEQ ID NO 30 for Pectinatus sp. DSM20764, SEQ ID NO 31 for Megasphaera cerevisiae, SEQ ID NO 32 for Selenomonas lacticifex, SEQ ID NO 33 for Zymophilus paucivorans and SEQ ID NO 34 for Zymophilus raffinosivorans.

As controls, the consensus probes SEQ ID NO 40 and 41 were used, which hybridise with the amplification products of all the species to be detected. Further possible binding sites for consensus probes are SEQ ID NO 42-45. The probes of SEQ ID NO 40 to 45 were determined by sequence comparison of known 23S rDNA and 5S rDNA sequences (GenBank Sequence Database, NCBI).

If the extinction measured for a 1 fg quantity of genomic DNA used in the PCR was greater than 1, the result was assessed as positive. The results of the PCR-ELISA are presented in Table 3.

_	
<u>ම</u>	
0	
۵	

			1,7			
	20	100 150 200 250	50 100 150 200 250 300	50 100 150 200 250 300 350	50 100 150 200 250 300 350 400	
		TCGAGGACTT AATATCTAGT CTGAAGGATA CGCCGATAGT	GTCGATAGGT TAATCAGTCG AACAGAGAAG ATGAAAAATA CCCATGCCGA GATCGCCCCC	G GTAGATAGGC C TAATCGGTCG A ACCGGAGCAT A TTCAGGGTTC A AATACACTGG A GAGCGCAAAG	G GTAGATAGGC IC TAATCGGTCG IA AACCGGAGCA IC GCTCAGGGTC IG ACAAGTACGT IG AGAGCGCAAA IC TGTTCCCATG IT GGTGGGAAAC	
	CAGGTAGATA		AGATGATCAG GGACTAATAC GCCAAGGTTG GCTCAGGCTT A TACACCTGTT	a AGATGATCAG C GGACCAGTAC A GCGCTTAGAA T GGCCAATGGA C GTTTCGATGA G CCAGTTTTGA	G AGATGATCAG G GGACCAGTAGA TT TGGCCATTGC TA CGTTTCGATG TA GGTAGTTTTG TA GGGATACAC TA GGGATACAC TA GGGATACAC TA GGGATACACC TA GAGATACACC TA GGGATACACC	
Sequence	上をひ上ならなられ	GAGGACTOR AGCGGACAGA TGTTTCGAGA TATAGTGTGG ACAGAAGTTA	GACTCCTGAA TATGTGAAGC ACAGGGTTAA AACGAAGTTC GCCTGAAGGA CACGCCAAAA	GACCCCTGAG TGCATGGAGC GTGAGCAGGA GCCGGGTTTT CTGCGAACGC A ACAATGATAG	A GACCCCTGAG A TGCATGGAGC G TGTGAGCAGG T GGCCGGGCTT G ACTGCGAACA A AACAATGATA C GATAGCAAGA T TCTTCACGCC	
		TAAGACCCCI TGAGGCGTGG ACAACGTAGT GAAGTTCTCT CATGCCGAAC	ATGGAAGTAA AGCATAGTGA CAAGGAAGAC GTTTTGAGAG GGTGGCGATA TAAGCTTCAG	ATGGAAGTAA AGTGCAGCGA CAAGTAGAGC GAGTTCGTTG GTTTCTGCGA CACAAAAACA	ATGGAAGTAA A AGTGCAGCGA C CAAGTAAGAG C TGAGCGTGAT A GGTTTCTGCG G GATAAGCT G GATAAGCT G GATA-3'	
		TATATGGAAG AGCAGCGCG AACCAAGTCA TTTGAGGGAA CACCTGTTCC AGTTGGGGGA			CCATTCCTAT TGGAAGTGGA AGGACTTAAC TAAGCGGGCC CTTATGTGCA TAAGTTCAAG GTTCTCATAA CCGAACACAG	
		່ເ	5, -	i ,s	2 2 2	┨
Description		23S-spacer-5S	23S-spacer-5S	23S-spacer-5S	23S-spacer-5S operon 1	
	Strain	DSM 20054	DSM 20690	DSM 20011	DSM 20008	-
Source	Species	brevis	lindneri	case.	paracasei ssp. paracasei	
nos	Genus	Lactobacillus	Lactobacillus	Lactobacillus	Lactobacillus	
SEQ ID	2 2	1	2	m	44	

						00000
	0 0	150 200 250 300	50 100 150 200 250 300	50 100 150 200 250 300	50 100 150 200 250 300	50 100 150 200 250 300
		TAATCGGTCG 1 TGCGACTGCG 1 ATTAAACGAT 2 TGGCGATAGC 2 AGCTTCTTCA 3	CCTGAGAGAT TGGAGCGGAC GTAGTGTTAG ACGTTCTCAG CCCATGTCGA GAGCACCCC	CCTGAGAGAT TGGAGCGGAC GTGGTGTTAG ACGTTCTCAG CCCATGTCGA GAGCACCCC	GACCCCTGAA TGTATGGAGC ATGTTAGGCT CGAAGTTCTT GAAGGATACA CCGATAGTAG	AGAGATGATC GCGGACCAAT AGAGAAGGAT CGAGCCGCGT ACACCTGTTC TAGTTGGGGG
		GGACCAGTAC TX AGCAGGTTTC TC CAAGGCAGCA A: ATAAGTGTGG TV ACAGAAGTTA ACAGAAGTTA ACAGAAGTTA A	AAGTAAGACC CCGTGAGGCA TAGCATGTAC GTGAGGCCA ATGAGGCGCA ATACACCTGTT CGTAGTTGGGG GTAGTTGGGG G	AAGTAAGACC C CCGTGAGGCA 1 TAGCATGTAC C TGAGAGCGCA A TACACCTGTT (GTAGTTGGGG (ATGGAAGTAA AGTACAGCGA CAAAGGTGCA TTTGAGAGAA GCGATAGCAA CTTCTTAGCG	AAGACCCCTG GATACATGGA GGTGTTCTCA TTCTTTCACA TGAGAAGGAT ACGCCGAGAG
Sequence			TCCTTTATGG AGTGGACGTG CTTAACCAAG TATCCAGTTT GCAAGAAGGA AGCGCCGAGA	TCCTTTATGG AGTGGACGTG CTTAACCAAG TATCCAGTTT GCAAGAAGGA AGCGCCGAGA	CCATTCCTTT TAGGAGTGGA AGGACTTAAC ATTATGCAGT TAGTGTGGTG AGAGTTAAG	TTATGGAAGT GAAGTGTAGT A ACCACAAAGT GAGAATAAAT GGGAATAAAT GTGACGATAGT F AAGCTTCTTA
	ATGGAAGTAA C		GATTTCCCAT ATAGGTTGGA CGGTCGAGGA AAGAAATGAA GGTGGCGATA TAAGCTTCTT	GATTTCCCAT ATAGGTTGGA CGGTCGAGGA AAGAAATGAA GGTGGCGATA TAAGCTTCTT	ATGAGATTTC GTAGATAGGC TAATCGGTCG AATATTACTT AAGCACAAAA TGTCGAACACAAAA ATGTCGAACACAAAA	T CCCATTCCAT G GTTGGGAGTG T CGAGGACTTA T TTAGTTTTGA G GAGAAGTGTG A CACAGAAGTTT T GCGAGGATAG
	上でいるよう		CTCGAGTTGA GATCAGGTAG CAATACTAAT TTTAAGGGCA AAAGTGGTGT ACACAGAAGT	- CTCGAGATGA GATCAGGTAG CAATACTAAT TTTAAGGGCA AAAGTGGTGT ACACAGAAGT	- ACGCCTCGAG AGATGATCAG GGACTAGTAC TTTGAAATGA CTCAGTGCGC CCTGTTCCCA	
		າ ກ	<u>v</u>	ហែ	- '2	ω ·
10:14:20	חפפרו	23S-spacer-5S operon 2	23S-spacer-5S	23S-spacer-5S	23S-spacer-5S	23S-spacer-5S
		DSM 20008	DSM 20001	DSM 20004	DSM 20019	DSM 20331
Source	urce	paracasei ssp. paracasei	coryniformis ssp. coryniformis	coryniformis ssp. torquens	curvatus	damnosus
	SOI	Lactobacillus p	Lactobacillus	Lactobacillus	Lactobacillus	Pediococcus
	SEQ ID	Sa lu	Φ	7	ω	O

			•		19		
	50	150 200 300 300	50	150 200 250 300 350 400	50	150 150 250 300 350 400 450	
Sequence	TAAGACCCCT GAGAGATGAT	GGTTGGGAGT GGAAGTGTAG TGATACATGG AGCGGACCAA TCGAGGACTT AACCACAAAG TGGTGTTCTC AAAGAGAAGA ATTTAGTTTT GAGAGAATAA ATTTCTTTCA CACGAGCGC CGGAGAAGTG TGGTGACGAT AGTGAGAAGG ATACACCTGT AACACAGAAG TTAAGCTTCT TAACGCCGAG AGTAGTTGGG	CGTGAAACCT GCCTTAAGAT GAGGTTTCCC	GGCTTGGAAG GCACCTTGAA TAAGACGAGG TAGATAGCCGA GTACAGTAAT GTACGAAGG GACTGGTACT AATAAGCCGA TAAAATCATC GAAAAAATG TTTGGTCTGA GATTTCTTCT GAGTGTGCAA GACACTCTGG TTGAAGGGCA GGGAACGTGA CTGCGGACTT TGGCTCAAAG AGTTAAAGCA TCTGGTGACG GGATCCACCT GTTCCCATTC CGAACACAGT AGTTAAGCAT AAGGTACTTG GGGGCGACC CCCTGGGAAA ATAGGACACT	GAGGTTTCCC	TTTGGAAGGC ACCTTGAAGA AGACGAGGTA GAIAGGCGG ATGGTGACAT ATGAAGCGGA CTGGTACTAA TAAGCCGAGA ATTTCATCAA AAAAGAGAAA TGTTTGGTCA GAGATTTTCT TTTGAGTGTG CAAGAACACT CGAGAGTATA TAGGTAAAGG GATAAGTTTC CTGGTTACTG TATATACCGG CTGAGGTGCT AGGCCAGAAC ATCTGGTGGC GATACCTGGA TGGATCCACC CCGAACACAG TAGTTAAGCA TCCACAGGCC GAAGGTACTT CCCCTGCGAA AATAGGACAC CGCC-3'	
	200	4 4 1 1 9 1 9 9 9 9 9 9 9 9 9 9 9 9 9 9	-\ <u>-</u>	<u> </u>			
Description		23S-spacer-55 9	22G_enacer-5S 5	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	30	23S-spacer-55	
		DSM 20285	200	DSM 20467		DSM 6306	
000	SOTTOS	inopinatus 2		cerevisiiphilus 2		frisingensis	
		Pediococcus		Pectinatus		Pectinatus	
	SEQ ID	01		11		12	

	1100 1150 2250 3350 4450 5500 5500	100 100 1150 2250 3300 350 400 450	50 100 150 200 250 350 350 450 500
2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	GCCTTAAGAT GAGGTITCCC GATGACGAG TAGATAGGCC GACTGGTACT AATAAGCCGA TGGTCCTGAT TTCTTCTGTG AGTATATAGG TAAAGGGAAA TATGAGCACT AAGGTGCAGA AAACTCACTT TGCGTGCTGA TTAGAGGAAA CGCGCGTTCA TTAGAGGAAA CACGCGTTCA TTAGAGGAAA CACGCGTTCA TTAGAGGAAA CACTGGATGG TGTGCGGAA CCTGGATGG	T GCCTTAAGAT GAGGTTTCCC A GATGACGAGG TAGATAGGCC GACTGGTACT AATAAGCCGA T TGTCCTGAT TTCTTCTGTG T GAGCACTAAG GTAAGGGGAA T GAGCACTAAG GTACTTCATC G CTCACTCTGC GTACTTCATC G CATCTGGTGG CGATACCTGG A GTAGTTAAGC ATCCACAGGC A AAGTAGGACA CCGCC-3	G AGGTTTCTCA TTACGAAAGT A ATAGGCCGGG AGTGGACGTA AT AGACCGAGA CTTGACTTAA G ACGCATAAAA TGGAGTGAGT AG CAGCGGAGA TACCAATGCA CGATTCGGAGT GGGTGAGGGA CGATTCGGAGT TACCAATGCA AC GACGAAAGAC TGATAAGATC AT ACGAATCCTG AAACGAATTC AT ACGAATCCTG AAACGAATTC AT CCCATACGA ACACAG-3' TT CCCATACGA ACACAG-3'
Sequence	AAGTGCTGAA AGCATCTAAG CGTGAAACCT AGAGCCGTAA GGCTTGGAAG GCACCTTGAA GGGAGTAGAA GTATGGTGAC ATACGAAGCG GAGCTTAACT TAATTTCATC TATAAATGTT AAGTTTTGAG TGTGCAAGAT CACTCATGAA GCAGCAGATT AGTTCCTGGT TTACTTTATA AAAGAACGTT TGAGGAAACG CGGCGTTCGT TTATCTCAAT GCTAAAGCAT TAAGATAATT CTAGCGTTCA CTCTGCGTAC TTATTTTCTA GGCAAGGAAA GCGTCGTTC GCGATGCTCA GACTGCTAAA GGCAAGGAAA CGCGTCGTTC TGAGGTGCTCA GACTGCTAAA GGCAAGGAAA CGCGTCGTTC CTGAAGCATC GACTGCTAAA GGCAAGGAAC CGCGTCGTCC GGCAAGGCACCCCC CTGCGAGGGTG GGTACTTGGG GGGCAGCCCC CTGCGAGGAGT	5 AAGTGCTGAA AGCATCTAAG CGTGAAACCT AGAGCCGTAA GGCTTGGAAG GCACCTTGAA GGGAGTAGAA GTATGGTGAC ATACGAAGCG GAGCTTAACT TAATTTCATC TATAAATGTT AAGTTTTGAG TGTGCAAGAT CACTCATGAA GCAGATTAGT TCCTGGTTTA CTTTATATAT GCAGATTAGT TCCTGGTTTA CTTTATATAT GCAACGTCTAA GGAAACGCGG CGTTCGTAGA TCTAGACTCC TAAAGCAGTA AGATCTGAAG ATGGATCCAC TGGGGGGGCAG CCCCCTGCGAA CGAAGGTACT TGGGGGGGCAG CCCCCTGCGAA	SY-GCATCTAAGC GTGAAACCAG CCTAGAGATG AAGTAAGGTC CCATGAAGAC GACATGGTAG CAGTAATGTA TGGAGCGGAC CGGTACTAAT GCAGGAACC CATTTTAAAG AAGCGAAGCG CGCTTATACC GAATCGCAGA TTCGGTAAAG GCGCAACAC CAGTTAGCAT AACTAAGC GTTTCGTAGC AGCGTAGGCT AACCCAACCA TGGTTTGAAA AAGAGTACAT GCGAAGAAAC AACATACAT TTCGATGTAG TTCTAGAAAC AACATACAA TTCGATGTAG TTGTCAGGAT AGTGGTGATG GCTGCAGGGA TCCACCTGTT
Description	23S-spacer-5S 5'-operon 1	23S-spacer-5S S operon 2	23S-spacer-5S operon 1
	DSM 20764	DSM 20764	DSM 20462
Source	zp.	വ ജ	a cerevisiae
	Pectinatus	Pectinatus	Megasphaera
OEO TD	13 13	14	15

	50	100 150 200 250 300	50 1100 1150 2200 250 3300 350 400	50 100 150 200 250 300	50 100 150 250 300 350
ı		AGTGGACGTA 1 CTTGACTTAA 1 GTTAGACGGG 2 TACGAATCCT 3 CCCATACGA 1	GAAGTATCTC GATAGGTTGG TAAATCGAGG GGTAACAGGT AGATGATTGG TTTTTCTTCT GTGACGATAG	F GACGICGCC F GAAGTATCTC A GATAGGITGG A TAAAICGAGG C TTCTGTATAG G ATAGCIGAGI C TCATACGCCG C TCATACGCCG C TCATACGCCG	T GAGGTTTCTC A GATAGGTCGG A TAGGTCGAGG IT TTCTTTCTTC TT CTGAGGCGAA G TGATAGCCAA G TGATAGCCAA G TGATGACGT AG TGCCC-3
	AGGTTTCTCA			T GCGAGAATAG T GTCCCGAGAT A AGACAAGGTA A CCAATACTAA T ACAATTTTTC A TCCAGTGACG TT AGTTAAGCACG	
Sequence	GLABABATA		CGTGAAGCCT CGTGAAGCCT CCTTGAAGA CCCAAGAAGCG A GAATCGAGTC A TCTAGAATGT CGTTCATTCA CCATACCGAA		
	- 1	GTGAAACCAG CCATGAAGAC TGGAGCGGAC AATAGAAAGA CCGAAATACT CAGTGGTGAT	AGCATCTAGG CCAGTAAGGT ATCGTAAGGT AACATCCCAA AACATCCCAA CCGCTAATAA CCGCTAATAA		
		5 · - GCATCTAACC AAGTAAGGTC CAGTAATGTA GCAAAGAAGC TAGTTAAGGT GAAACGAATT ACACAG-3 ·	- AAGTGCTGAA ATGGAGTAAT GAGTGTAACTTT GCTTAACTTT CGTATGCGAA CAGATGTTGA GTATAGTTTTA	ACGCCGAAGG -3. 5 AAGTGCTGAA ATGGAGTAAT GAGTGTAAT GCTTATCTTA TTTTGAGTGG GGTACCCT	AAAGTACTTGAA ACAGAGCAAT GAGTGGAAGC GCTTGACTTA TGTATAGTTT AGCGAAGGC GTGGAAGGC CGAAAGTACT CGAAAGTACT
		- , 5	<u></u>		S S
-	Description 	23S-spacer-5S operon 2	23S-spacer-5S operon 1	23S-spacer-5S operon 2	23S-spacer-5S
		DSM 2	DSM 20757	DSM 20757	DSM 20765
	Source	Φ	lacticifex	lacticifex	1S raffinosivorans
		Megasphaera cerevisia	Selenomonas	Selenomonas	Zymophilus
	SEQ ID	NO 16	17	18	19



	50	100 150 250 300 350	
Sequence	Description Corps ACCTTAAGAT GAGGTTTCTC 50	5 AAGTGCTGAA AGCALCIANO CCCTTGAAGA AGACAAGGTA GATAGGTCGG 1100 ACAGAGCAAT CTGGTAAGAC CCCTTGAGA CCGATACTAA TAGGTCGAGG 150 GAGTGGAAGC GCAGTAATGT GTGTAGGAACAT TTCTTTCTTC 200 GCTTGACTTA AAGCCAGAAC GAATTCTAAG ATGCGAACAT TTCTTTCTTC 200 TGTATAGTTT TGAGAGAACA GACTCTTAAG ATGAGCAGTC TGAGGCGAAA 250 TGTATAGTTT TGAGAGAACA AAAAAAGAA TATCTGGTAG TGATAGCCAA 3100 GCTAAAGGCA GCGATATCTA AAAAAAGAA TATCTGGTAG ACTTGAACCAA 3100 GTGGACCCAC CTGTTCCCAT ACCGAACACA GTAGTTAAGC ACTTGAACGT 350 GTGGAAGTACT TGGGTGGAAA CGCCCTGGGA AAATAGGACA CCGCC-3.	
	Description	23S-spacer-5S	
		DSM 20759	
	Source	Zymophilus paucivorans	
		Zymophilus	
	SEQ ID	NO 2 0 0	

8	
<u>o</u>	
ab	
Ë	

	-3,	-31	,	- '	7	-3,	-3,	-3,	-31	-3,	-3,	-3,	-3,	-3,	2 6	<u> </u>	-3	7	2	-3	1.5	<u>,</u>	-3	-
Sequence	הטורות גוונט גייט יייט יי	CCAAGTCAACAAGTAGT	GACACAGGGTTAAATCAAAGTTG		AGGTTTCTGCGACTGCGAAC	ATGTACGTAGTGTTTAAGGGCC	CHINCHCAGGCACA	CITCLCAGAGAAGGATTCG	GTTCTCAAAGAGAAGATTTCGATATTA	TGAGAGCGTAAAACTGCGGACTT	CAGATAAGTTTCCTGGTTACTG	CACTAAGGTGCAGAAAAGAACGT	CTTTTCGATGTAGTTGTCAGGATACG	GTTCATTCAATAATATCCAGTGACG	AACTCTTAAGATGGAGYAGTCTG	ACTCTTAAGATGAGCAGTCTGA	The standard of the standard o	ACTOLOGICA	GTGAAGTTTTGAGTGTGCAAGA	GACCGAGGACTTGACTTAAGCA		TCCAGTGACGATAGCTGAGT	AAGAATATCTGGTAGTGATAGCCAA	
		51-	51-		5,-	5,-		5,-	- 10	- , 5		<u>'</u> '		2,5	, 4		,	5,-	51-	1	,	5,2	1	
		Specific probe	probe		Specific probe	Specific probe		Specific probe	Specific probe	Specific probe	Specific probe	Specific probe	Specific probe	Specific probe	Specific probe	Specific probe	Specific prope	genus-spectration	genus-specific	genus-specific	probe	genus-specific	genus-specific	probe
-	Description	- 11	brevis	lindnerı	casei paracasei ssp. paracasei		Lactobacillus coryniformis ssp.torquens		1	inopinatus	cerevisiiphilus	frisingensis	sp. DSM 20764	cerevisiae	lacticifex	raffinosivorans	paucivorans	genns	genus	pliner	anilo 6	genus	255	Spireh
			Lactobacillus	Lactobacillus lindner	Lactobacillus Lactobacillus	Lactobacillus	Lactobacillus	Sullinedotori	pediococcus	Pediococcus	Pectinatus	Pectinatus	Pectinatus	Megasphaera	Selenomonas	Zymophilus	Zymophilus	Pediococcus	Pectinatus		Megasphaera 	Selenomonas		Zymophilus
	SEQ ID	NO	21 1	22		24		2		27	α,ς	0 0	30	31	32	33	34	35	36		37	38		39

_
آن
=
0
Ŭ,
=
2
ø
≗
ڡؚ
a
⊢

	-3,	-31	-3,	-31	-31	-31	-3,	18-	-3	-31	-3,	-3,	-31	-3,	15-	, (16-	-3,		-3.	-3,	-3,	-3,	-3,	-31	-3,	-31	-3,	-31	-3,	15-		, <u> </u>	
Sequence	び出立びと目出てよびよびよび	GTCGTGAGACAGI	CYTAGTACGAGGACCGGRA	GCTACCTGGGGATAACAGGC	ATCGACGGGAGGTTTSSCAC	CACCTCGATGTCGGCTCRIC	CCAAGGGTTGGGCTGTTC	AAGGCCATCRCTCAACGG	AAGTGCTGAAAGCATCTAAG	TGTGTTCGilATGGGAACAGGIG	TGTGTTCGGAATGGGAACAGTG	TGTGTTCGAAATGGGAACAGG IG	TGTGTTCGGTATGGGAACAGGTG	TGTGTTCGATATGGGGAACAGGIG	TGTGTTCGGCATGGGAACAGGTG	TGTGTTCGACATGGGAACAGGTG	GGCRRYGTCCTAYTYTCSC	GGCAGTGTCCTACTTTCCC	GGCAGCGTCCTACTTTCGC	GGCAGTGTCCTACTTTCGC	GCCTCCTACTTTCCC	COURTEMPORTING TO STATE OF THE	DOLLEGO TONICE ID	GCTTANCI ICCOLOR	GCTTRACTICIATOR	GCTTPACTICION COMMENT AND	GCTTAACTICCATOT	GCTTAACTTICCGGG11CG	GCTTAACTTCTAGGTTCG	GCTTAACTTCTGGGTTCG	GCTTAACTTCCAGGTTCG	GCTTAACTTCCGAGTTCG	GCTTAACTTCTAAGTTCG	
		- 15	51-	51-	5,-	5,-	5	51-	5,-	- 12	5	- 15	5 -	51-	51-	2,4	51-	5,-	- 12		5,-	5, -	5 ' -	5 -	5,-	51-	- , 5	51-	5,-	5	51-	- 15	5 - 2	
	Description		consensus sequence	consensus sequence		consensus sequence	consensus sequence	consensus sequence	consensus sequence	consensus sequence	consensus sequence	consensus sequence	consensus sequence	consensus sequence	consensus sequence	1	1	1			- 1	- 1			- 1	- 1	consensus sequence	consensus sequence	consensus sequence	consensus sequence	consensus sequence		1	
	SEQ ID	NO	40 00			T			T		T	49 C	50 C	Γ		1	T	T	Τ	7			59	09	61	62	63	64	65	99	67	89	69	

							(-1		[-]	-	-	1-	'n		25 T			3,	-3,	3,	3,1	3,1) T	3,	-3,	3,	-3,	-3,	- ? . -]	
	-3,	1	1		<u>~</u>	-3	-31	-3,		-3	-3,	-3,	-3,	-3,	1-31	4'	4'	+	-3,			-		-	1	AG.	ı				+	+	
Sequence	COMBONDE	GCTTAACTTCTGAGTTCG	GCTTAACTTCCAAGTTCG	TCGAGAATAATTGAATAATATCTAG	GAGGGAAGATCTCTTAT	A A A A A A A A A A A A A A A A TA TA TCTAGTT	AACAGAGAGAGAGAAAAAAAAAAAAAAAAAAAAAAAAAA	TTGAGAGAACGAAGTTCGCICAGGCIIATCTGTGTGTGGG	TTCGTTGGCCGGGTTTTGGCCAAIGGAIICAGGTTTCTTTGGTTTTTGGTTTTTTGGTTTTTTGGTTTTTT	GCGTTTCGATGAAATACACTGGTTCCCAGTT	SACCEPTA A SACCEPTA A GCG TGA	1 I I I I I I I I I I I I I I I I I I I	GCGTGATGCCCGGCTTTCCCATTAAACAATGATAGTT	CAAGTACGTTAGGT COMMINGAGAGGGGAACGTTCTCAGAAA	AAAGAAATGAATATCCAGIIIIGAGAGGGGGGGGAAAAAAATGCAGT	AGGTGCAATGTTAGGCTTTTGAAATGAAAIAITACITTTTTTTTTT	GCCGCGTAAGTGGATCGGAAA	GCCGCGGAAGTCGGAGAA	KCC KC RCHILLE	GAGAGAATAAATTTCTTTCACACGA	TOTAL STATE OF STATE OF THE STA	AAAATCATCAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	CACTICITED I GARGOCCIO CONTROLLO CON	GATTTCATCAAAAGAGAGAATGI I I COT CITCLE	TATATACCGGCTGAGGTGCTGAGGCACTCAGGGCTGCTTGAGGGTGGTGAGGGTGGTGGTGGGTG	AATTTCATCTATAAATGTTTGGTCCTCATTTTCAGAAAA	AGATTAGTTCCTGGTTTACTTTALATAGACCACTTTCCTTAGATTAGA	AGGAAACGCGGCGTTCGTAA	TAATAATCTAGAATGTTTCGATACAATTTTTTTTTCTGTATAGTTTTTGAG	TGGGCAGCAGCAT	GAGGCGAAAGCTAAAGGCAGCGAT	AATCCTGAAACGAATTCAGTGGTGATGGCTGCAGGGA	
		5'-	1,'	+	- 1	<u> </u>	- ' '	-1.9		51-		5,-	5,-	5 ' -	- , 5	51-	51.2	5 \ -	, _	5,-		5,2	- 15	5'-	5,-	- 12	5,-	<u>'</u>	7 [5	- ;	5 5	2 2	2
		5	1		probe		specific probe 5'	probe		nrobe		specific probe 5'-	specific probe 5'	specific probe 5'	specific probe 5'-			probe	specific prope	specific probe		specific probe	specific probe 5'	specific probe 5'		specific probe		specific probe		- 1	specific probe	specific probe 5'-	specific prop
- - - -	Description				brevis	hrevis	יישמליי	Tildiera	Inquer	Caseı	casei	casei	paracasei	i do a concrete	paraces	corynilotinia	curvatus	damnosus	inopinatus	damnosus	inopinatus samijus	gervarieiinhilus			rrisingenera	ITISIIIGEIBES	Sp. Dan 2010		sp. DSM 20764	lacticifex	raffinosivorans	paucivorans	grevisiae
			consensus seguence		lus					Lactobacillus	Lactobacillus					Lactobacillus	Lactobacillus	Pediococcus	Pediococcus	Pediococcus	Pediococcus	Pediococcus	Pectinatus	Pectinatus	Pectinatus	Pectinatus	Pectinatus	Pectinatus	Pectinatus	Selenomonas	zymonhi lus	Zymophilus	
	EQ ID	NO	71 C	T	T		74 I	75 I	1 9 <i>L</i>	77 I	78	79		80	81	82	83		85	86			87	88	89	90	91	92	93	94	100	200	3

Table 2 (Cont.)

_
٦
0
Ö
ン
2
Φ
亙
<u></u>
⊢

		Description		Sequence	
SEQ ID	1	4			,
NO			1	TATGGAAGTAAGACCCCTGA	. 2
c			' '	上しただな中でしたのでは、これでは、	-3,
βŔ	,		5,-	AGATGATCAGGTAGGTAGGGT	
100	Detection of all Lactobacillacea	Detection of all Lactobacillaceae relevant to brewing	5,-	AGATGATCAGGTCGATAGGTT	-3,
	for differentiation if	חוו סכווכד אמכסכאבי		EECC VE	13.
	brewing		51-	AGATGATCAGGTAGATAGGTI	,
101			, 4	TACTAATCGGTCGAGGACTTAACCA	-3,
5				ADDA ATTO ADDA ADDA ABO ATT	-3`
102			5,-	ATACTAGLOGROGICATION	10-
103		Canal Supering	5,-	GAAGCGGACTGGTACTAATAAGCCGAGAGCTT)
104					
1	Pectinatus	denna		A SOCIAL STATE OF STATE STATE SACRESCENTA	-3,
,		denn	2,2	CAGCGGACCAGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	
SOT	Selenomonas	genus		A A THIT A SHIPT SO SO SO SECOND SECO	-3,
		ific	5,-	AGCGGACCGATACTAGGICGAGGCIIGGCII	
106	Zvmophilus	genus		出土し 女がじ なびしつ なびな出な な出り な出からから エジニュニュー	-3,
i d	4	nueb	5,-	GGACGGACCGGIACIAATAGACCGGGGGG	
/07	Megasphaera	genus			

~
<u>a</u>
٥
ā
$\overline{}$

			OEO ID	SEO ID	SFO ID	SEQ ID SEQ ID SEQ ID SEQ ID	SEQ ID	SEQ ID	SEQ ID		\vdash	_	SEQ ID
		NO 21	NO 2	NO 23		NO 25	NO 26	NO 27	NO 28	NO 29	NO 30	NO 31	NO 32
		1	-		-		•	1	-	-	-	.	
Lactobacillus brevis	brevis	۲								1	1	1	•
Lactobacillus lindneri	lindneri		+		'								
Lactobacillus	casei	•	'	+									•
l actobacillus	paracasei paracasei	•	•	+	-	•							'
Lactobacillus coryniformis	coryniformis			•	+								
	coryniformis								'	'		•	ı
l actobacillus	actobacillus coryniformis torquens	•	•	•	+	•	•					•	
l actobacillus	Curvatus			•		+	•		'				
במכוססמכוומס				'			+	-	<u> </u>				
Pediococcus	damnosus							+	1	1		•	<u>'</u>
Pediococcus inopinatus	inopinatus		-		•				+		'	•	•
Pectinatus	cerevisiiphilus							.		+			•
Pectinatus	frisingensis					•				'	+		
Pectinatus	sp. DSM 20462	-		•	-				-			+	•
Megasphaera cerevisiae	cerevisiae	•		-	.					.		'	+
Selenomonas lacticifex	lacticifex	,		•	-			, ,	-	-		-	
Zymophilus	raffinosivorans	-		-	-	-			\\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	-			
Zymophilus	paucivorans				-	-							

		SFO ID	SEO ID	SEG ID SEG ID SEG ID SEG ID SEG ID SEG ID	SEQ ID				
		NO 33	NO 34	NO 35	NO 36	NO 37	NO 38	NO 39	NO 39 NO 40-45
l actobacillus brevis	revis				,		-	1	+
l actobacillus lindneri	ndneri				1		-	•	+
l actobacillus casei	igo				•	•	•	ı	+
l actobacillus p	actobacillus paracasei paracasei	-				-	•	•	+
Lactobacillus	Lactobacillus corvniformis			•	-	-	1	J	+
Lactobacillus	Lactobacillus coryniformis torquens			-	-	1	•	-	+
Lactobacillus curvatus	urvatus			•	-	1	1		+
Pediococcus damnosus	Jamnosus		,	+		1	•	•	+
Pediococcus inopinatus	nopinatus			+		•	-	•	+
Pectinatus	cerevisiiphilus			,	+	-	•	•	+
	frisingensis				+	1	•	•	+
	sp. DSM 20462		•		+	1	'	'	+
l a	cerevisiae	'	,	•	•	+	-	•	+
Selenomonas lacticifex	acticifex				•	,	+		+
Zymophilus raffinosivorans	raffinosivorans	+	•	-	'	•		+	+
Zymophilus	paucivorans		+	•	•	•		+	+